Brief description of the activities during the applicable regulatory review period. IND Activities

Date of	
Contact	Summary of Contact
ooneu o c	
3-Feb-1993	FDA Acknowledged receipt of IND submission (SN000, dated January 27, 1993).
16-Feb-1993	Response to request for information.
15-Mar-1993	Protocol amendment
2-Jul-1993	Request for meeting
10-Aug-1993	Comments from Agency and request for teleconference.
17-Aug-1993	Response to request for information.
24-Sep-1993	Minutes from clinical development plan meeting held on September 3, 1993. Revised meeting minutes from September 3, 1993 clinical
4-Nov-1993	development plan meeting reflecting the Agency's comments.
10-Dec-1993	Information amendment
14-Jan-1994	Protocol amendment
8-Apr-1994	Comments from FDA regarding IND amendment dated December 10, 1993
27-Apr-1994	IND annual report
20-May-1994	New protocol
18-Jul-1994	Protocol amendment
18-Oct-1994	Protocol amendment
4-Nov-1994	Letter of Authorization regarding IND 41,574.
6-Dec-1994	Regarding obtaining written guideline on interactive IND process and inquiring how much time new MRO requests to review draft Phase III protocols.
20-Jan-1995	Information amendment
3-Mar-1995	Request for designation; recommendation for primary review authority be given to Pilot Drug Division under jurisdiction of CDER.
13-Mar-1995	Acceptance of the request for designation (dated March 13, 1995)
24-Mar-1995	Request for an end of Phase II meeting
4-Apr-1995	EOP II meeting Briefing Package.
28-Apr-1995	Information amendment
3-May-1995	Annual report
9-May-1995	Response to request for status of requested CMC meeting with device and Pilot Drug reviewers. Regarding request for designation of CDER as the agency with
12-May-1995	primary jurisdiction for the pre-market review and regulation of the product.
22-May-1995	Date and time for CMC meeting with Pilot and CDRH set for June 27, 1995.

D. 5 - 5	
Date of	Summary of Contract
Contact	Summary of Contact
0 7 1005	Five draft pivotal protocols submitted for FDA review and
2-Jun-1995	comment.
1005	Inquiry regarding review status of five Phase III protocols
9-Jun-1995	(sent June 2, 1995); FDA will fax comments by June 15, 1995.
14-Jun-1995	Comments regarding submission dated April 28, 1994
	Draft meeting notes from the FDA/Janssen/ALZA meeting held
13-Jul-1995	June 20, 1995 to discuss CMC issues.
	Response FDA comments on five pivotal protocols (fax dated
1-Aug-1995	June 14, 1995).
14-Aug-1995	Teleconference on August 14, 1995 per ALZA request to
to 15-Aug-	discuss clarification of written comment concerning C-95-
1995	019.
17 7 1005	
17-Aug-1995	Protocol amendment
	Excerpted information from the summary basis of approval for
1005	LAAM and the MRO's overview of the "usage" study which is
28-Aug-1995	required for ETS fentanyl.
1 0 - 1005	Final meeting minutes from the June 27, 1995 meeting with
1-Sep-1995	Pilot Drug and CDRH representatives to discuss CMC issues.
1 0 1005	Draft meeting notes from June 27, 1995 CMC meeting have been reviewed without comment from FDA. Request for information
1-Sep-1995	
	Response to call from FDA regarding final minutes from the
01 0 1005	June 27, 1995 meeting. Request for information requested at
21-Sep-1995	the meeting. ALZA response.
	Draft qualification plan in follow-up to the June 27, 1995
21-Sep-1995	meeting with ALZA/Janssen/FDA.
	Response to questions from Dr. Lee about the draft
	Qualification Plan (dated September 21, 1995) asking about
20 000 1005	electrical current density and dermal responses after
29-Sep-1995	administration of ETS (fentanyl).
10 Oct 1005	Request for late November or early December End of Phase II
19-Oct-1995	meeting. Protocol amendment - new protocol C-95-039; information
27 0-+ 1005	amendment - CMC.
27-Oct-1995	Confirmed date of November 28, 1995 for End of Phase II
0 Nov. 1005	·
8-Nov-1995	meeting. Pre-meeting package for November 28, 1995 End of Phase II
13-Nov-1995	meeting.
13-100-1993	Protocol amendment - new protocol C-95-032, amendment to C-
17-Nov-1995	95-019; information amendment - CMC.
17-100-1995	Request for clarification of response to ALZA's proposed ETS
7-Dec-1995	placebo design at End of Phase II meeting.
7-Dec-1995	Response to ALZA's December 20, 1995 phone request for
21-Dec-1995	information on the closed session of the Advisory Committee.
21-DeC-1993	Meeting minutes from End of Phase II meeting held on
22-Dec-1995	November 28, 1995.
22-DEC-1993	Inquiry about status of the End of Phase II meeting minutes
	(sent December 22, 1995) and date for closed session of Life
16-Jan-1996	Support and Anesthetics Advisory Committee.
10-0411-1330	Confirm April 30, 1996 Advisory Committee meeting and see if
	there was a final agenda for allotted time. Inquiry about
	status of draft EOP2 meeting minutes (sent December 22,
15-Fob-1996	1995). Re-confirmed acceptability of C-95-023.
15-Feb-1996	1133). Re-Contitumed acceptability of C-33-023.

D-to of	
Date of	Summary of Contact
Contact	
8-Feb-1996	Preclinical and clinical topical safety data related to inquiry from June 27, 1995 meeting.
0-LED-1930	
11 1 1 1000	Agenda for April 30, 1996 closed session of Anesthetics and
11-Mar-1996	Life Support Advisory Committee meeting.
	Inquiring if the proposed plan not to perform ex-US pivotal
	trials under the IND would create any problems at FDA in
	terms of the interactive IND. FDA request for outline
14-Mar-1996	labeling. ALZA faxed response.
1005	Copy of the abbreviated draft labeling, provided per FDA
14-Mar-1996	request.
	FDA issues for April 30, 1996 closed session of Anesthetics
18-Mar-1996	and Life Support Advisory Committee meeting.
	Draft package for April 30, 1996 closed session of
28-Mar-1996	Anesthetics and Life Support Advisory Committee meeting.
•	Final package for April 30, 1996 closed session of
	Anesthetics and Life Support Advisory Committee meeting.
5-Apr-1996	Same as March 28, 1996 package to Dr. Burke.
16-Apr-1996	Corrected FDA sheets for Advisory Committee. ALZA faxed
to 17 Apr,	response of corrections and agenda suggestions. ALZA call to
1996	discuss meeting logistics.
	Draft agenda for April 30, 1996 closed session Advisory
17-Apr-1996	Committee meeting with ALZA/Janssen/FDA per phone request.
3-Jun-1996	Comments from the April 30, 1996 Advisory Committee Meeting.
	Protocol amendment - new protocol C-96-020; information
7-May-1996	amendment - CMC.
	Annual report covering the period of February 27, 1995 to
14-May-1996	February 26, 1996.
	Minutes from April 30, 1996 Closed Advisory Committee
	Meeting with Division of Anesthetics, Critical Care, and
10-May-1996	Addiction Drug Products.
	Protocol amendment - new protocol C-96-007; information
20-May-1996	amendment - CMC.
	Protocol amendment - new protocol C-96-029; CMC update; new
3-Jul-1996	investigator
	Protocol amendment - change in protocol C-96-020 (reference
3-Jul-1996	SN022, dated May 7, 1996).
	Faxed info for teleconference regarding definition of
	clinically relevant respiratory depression in trials. FDA
	would like to base definition on respiratory rate and
9-Jul-1996	sedation only.
7-Aug-1996	August 7, 1996 phone call indicating FDA would like a
to 8-Aug-	teleconference to discuss some issues on the clinical
1996	program. ALZA faxed response dated August 8, 1996.
	Protocol amendment - new protocols C-96-006 and C-96-009;
16-Aug-1996	CMC update.
	Teleconference to discuss FDA comments (FDA fax dated August
	8, 1996) on Phase I protocols for JAN-2, and to discuss
	specific issues in relation to JAN-1 (ALZA fax dated August
22-Aug-1996_	8, 1996).
	Response to device questions raised at the April 30, 1996
11-Sep-1996	Life Support and Anesthetics Advisory Committee.
17-Sep-1996	Protocol amendment - change in protocol C-96-006.
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Date of	
Contact	Summary of Contact
COIICACC	Dummary Of Contract
24-Sep-1996	Protocol amendment - change in protocol C-96-009.
	Pharmacokinetics reviewer comments on skin tolerance
29-Sep-1996	protocol C-96-029 (submitted in SN025, dated July 3, 1996).
	Teleconference to discuss a question on recent amendment to
	protocol C-96-006. Comments faxed after teleconference
3-Oct-1996	concerning study C-96-009.
	Response to FDA comments (fax dated September 29, 1996) on
'	protocol C-96-029-02 (SN25, dated July 3, 1996) regarding
15-Oct-1996	calculation of amount of fentanyl delivered.
	Response to FDA comments faxed on October 3, 1996 regarding
25-Oct-1996	Phase I protocol C-96-009 (SN027, dated August 16, 1996).
	Position paper regarding difficulty in distinguishing
	topical effects related to electrical current versus
	chemical effects related to the delivery of fentanyl from E-
20-Nov-1996	TRANS (fentanyl) systems.
	Phase III randomized controlled studies; design and
26-Nov-1996	statistical analysis features submitted for FDA comment.
20 1101 1330	Pharmacokinetic comments from FDA regarding study C-96-009
2-Dec-1996	submitted in SN027 (dated August 19, 1996).
, 500 1550	FDA comments regarding the definition of "clinically
	relevant respiratory depression" in protocols C-97-058-02,
27-Dec-1996	C-94-059-02, and C-95-016-02.
27 BEC 1330	C 94 039 02, and C 93 010 02.
6-Jan-1997	Protocol amendment - new protocol C-95-016; CMC update.
	Minutes of conference call held between Division of
	Anesthetic, Critical Care and Addiction Drug
6-Jan-1997	Products/ALZA/Janssen.
	Request for clinical studies filed to INDs 41,574 and
	50,284. ALZA request for clarification of FDA request dated
9-Jan-1997	January 2, 1997
	Information amendment - clinical with respect to study C-95-
17-Jan-1997	016
17-Jan-1997	January 17, 1997 request for clarification of FDA phone
to 21-Jan-	request dated January 9, 1997. January 21, 1997 FDA phone
1997	response.
	Response to request for clarification on which clinical
	protocols were filed to IND 41,574 and IND 50,284 and
23-Jan-1997	request for clinical development plans for both projects.
	Draft risk analysis standard operating procedure and interim
31-Jan-1997	risk analysis.
	Proposed finished product specification and rationale
10-Feb-1997	documentation; follow-up meeting request.
	General correspondence - Submitting 3 Phase III protocols
19-Feb-1997	utilizing a reduced on-demand dosage for comment.
	Statistician's comments on Phase III pivotal protocols C-94-
12-Mar-1997	057-03, C-94-058-02, C-94-059-02, and C-95-016-02
	Medical Officer completed review of the 25 µg Phase III
	protocols (SNO40, dated February 19, 1997). Ruled to be safe
31-Mar-1997	to proceed.
	Information amendment - toxicology; general correspondence -
3-Apr-1997	meeting request.
	Response to request for information on INDs 41,574 and
3-Apr-1997	50,284
<u> </u>	100/100

Date of Contact	Summary of Contact
11-Apr-1997	Request that we assign serial numbers and FDA Form 1571s for the April 3, 1997 submission of CMC material
14-Apr-1997	Form 1571s to complete April 3, 1997 submission of CMC information
23-Apr-1997	Response to request for copy of ALZA regulatory standard (Code #0007075).
11-Apr-1997	Copy of the transcripts from the Closed session of the April 30, 1996 Anesthetic and Life Support Advisory Committee.
18-Apr-1997	Response to statistical comments on Phase III protocols.
2-May-1997	Request for more detailed information on the primary package container materials, which report to justify CPC lower limit on the final product specification, and to change wording for the CPC specification.
8-May-1997	Response to phone request from FDA regarding CMC information.
8-May-1997	Annual report covering the period of February 27, 1996 to February 26, 1997.
16-May-1997	General correspondence - Phase III protocol C-96-057. Forwarding FDA's comments regarding Interim Risk Analysis
17-May-1997	(SN038, dated January 31, 1997).
17-May-1997	Contact information the new CSO for the project.
30-Jun-1997	Request for meeting on CMC information requirements. Follow-up to last CMC meeting request (dated June 30, 1997),
14-Aug-1997	ALZA ready to submit package, waiting for meeting date.
15-Aug-1997	Background package for meeting on CMC. Response to Medical Review Officer's comments on interim
29-Aug-1997	risk analysis performed.
29-Aug-1997	Protocol amendment - change in protocol for study C-95-016. FDA internal meeting on CMC package planned for September 10, 1997, postponed due to more work needed. Drug
9-Sep-1997	development held up. Minutes from FDA teleconference dated September 25, 1997 to
29-Sep-1997	discuss registration batch plans.
28-Oct-1997	Protocol Amendment - new protocol; C-97-001; CMC update. Protocol amendment - change to Phase I protocol C-97-001
7-Nov-1997 12-Nov-1997	(FEN-USA-63). Response to CMC questions received via fax on August 21, 1997 in relation to original IND (SN000), SN039 (dated February 10, 1997), SN041 (dated April 3, 1997), and SN044 (dated May 8, 1997). Protocol amendment - new protocol FEN-USA-29 (C-94-057), new
24-Nov-1997	investigator
5-Dec-1997	Protocol amendment - new protocol FEN-USA-28 (C-94-060), new investigator
5-Dec-1997	Protocol amendment - new protocol FEN-USA-58 (C-96-055), new investigator
5-Dec-1997	Response to Pharmacology questions received via fax on August 21, 1997 from CSO in relation to SNO41 (dated April 3, 1997).

Date of Contact	Summary of Contact
Odiredo	Follow-up on request for meeting with CDRH regarding device
	content of the NDA, timing of pre-NDA meeting, and label
15-Dec-1997	utilization study.
	Call regarding the IND amendment for PK study C-97-001. No
	issues with the study, but would like to know the status of
18-Dec-1997	the study.
	FDA suggested information for pre-NDA package in response to
14-Jan-1998	ALZA queries.
14-Jan-1998	Protocol amendment - new protocol C-96-056.
11 0411 1990	Reviewer wanted to know if C-94-057 was a pivotal study.
20-Jan-1998	ALZA responded that it was.
20-Jan-1998	Protocol amendment - new protocol C-94-068.
	Follow-up on phone messages regarding pivotal status of study C-94-057 (dated January 20, 1998). Faxed reference to
122 Tan-1000	where documents have been filed.
23-Jan-1996.	Protocol amendment - new investigator documents for studies
2-Feb-1998	FEN-USA-29 and FEN-USA-58.
Z-FED-1990	
3-Feb-1998	Protocol amendment - change to Phase I protocol C-94-060.
9-Feb-1998	Protocol amendment - new protocol C-96-057.
J 16D 1550	Protocol amendment - new investigator documentation for C-
27-Feb-1998	196-055.
27 102 1330	Protocol amendment - new investigator for study FEN-USA-28
3-Mar-1998	(C-94-060).
9-Mar-1998	Initial IND safety report (Ref. #C000201).
1000	Completed review of SN054 (dated November 25, 1997) and
13-Mar-1998	SN056 (dated December 5, 1997); FDA requests and comments.
	Voicemail acknowledging receipt of request to meet with CDER and CDRH regarding device documentation proposal for the
24-Mar-1998	NDA.
24-Mai-1990	Stating that our formal request for a CDER/CDRH
	teleconference to discuss our proposal for device
25-Mar-1998	
	content/format of the NDA has been granted.
	content/format of the NDA has been granted. Response to FDA fax regarding statistical comments on
22-Apr-1998	content/format of the NDA has been granted.
	content/format of the NDA has been granted. Response to FDA fax regarding statistical comments on protocol C-94-057 and protocol FEN-USA-58.
22-Apr-1998	content/format of the NDA has been granted. Response to FDA fax regarding statistical comments on protocol C-94-057 and protocol FEN-USA-58. Notification of temporary suspension of clinical trials due
22-Apr-1998	content/format of the NDA has been granted. Response to FDA fax regarding statistical comments on protocol C-94-057 and protocol FEN-USA-58. Notification of temporary suspension of clinical trials due to non-safety related technical issues.
22-Apr-1998	content/format of the NDA has been granted. Response to FDA fax regarding statistical comments on protocol C-94-057 and protocol FEN-USA-58. Notification of temporary suspension of clinical trials due to non-safety related technical issues. Pre-meeting package for intercenter teleconference with CDER and CDRH representatives to discuss the device related aspects of the NDA.
22-Apr-1998 29-Apr-1998 11-May-1998	content/format of the NDA has been granted. Response to FDA fax regarding statistical comments on protocol C-94-057 and protocol FEN-USA-58. Notification of temporary suspension of clinical trials due to non-safety related technical issues. Pre-meeting package for intercenter teleconference with CDER and CDRH representatives to discuss the device related aspects of the NDA. Verification of receipt of three copies of pre-meeting
22-Apr-1998 29-Apr-1998	content/format of the NDA has been granted. Response to FDA fax regarding statistical comments on protocol C-94-057 and protocol FEN-USA-58. Notification of temporary suspension of clinical trials due to non-safety related technical issues. Pre-meeting package for intercenter teleconference with CDER and CDRH representatives to discuss the device related aspects of the NDA. Verification of receipt of three copies of pre-meeting package; would like additional five copies.
22-Apr-1998 29-Apr-1998 11-May-1998 15-May-1998	content/format of the NDA has been granted. Response to FDA fax regarding statistical comments on protocol C-94-057 and protocol FEN-USA-58. Notification of temporary suspension of clinical trials due to non-safety related technical issues. Pre-meeting package for intercenter teleconference with CDER and CDRH representatives to discuss the device related aspects of the NDA. Verification of receipt of three copies of pre-meeting package; would like additional five copies. Response to request for five additional copies of SNO69
22-Apr-1998 29-Apr-1998 11-May-1998	content/format of the NDA has been granted. Response to FDA fax regarding statistical comments on protocol C-94-057 and protocol FEN-USA-58. Notification of temporary suspension of clinical trials due to non-safety related technical issues. Pre-meeting package for intercenter teleconference with CDER and CDRH representatives to discuss the device related aspects of the NDA. Verification of receipt of three copies of pre-meeting package; would like additional five copies. Response to request for five additional copies of SNO69 (dated May 11, 1998).
22-Apr-1998 29-Apr-1998 11-May-1998 15-May-1998	content/format of the NDA has been granted. Response to FDA fax regarding statistical comments on protocol C-94-057 and protocol FEN-USA-58. Notification of temporary suspension of clinical trials due to non-safety related technical issues. Pre-meeting package for intercenter teleconference with CDER and CDRH representatives to discuss the device related aspects of the NDA. Verification of receipt of three copies of pre-meeting package; would like additional five copies. Response to request for five additional copies of SN069 (dated May 11, 1998). Annual report covering the period of February 27, 1997 to
22-Apr-1998 29-Apr-1998 11-May-1998 15-May-1998	content/format of the NDA has been granted. Response to FDA fax regarding statistical comments on protocol C-94-057 and protocol FEN-USA-58. Notification of temporary suspension of clinical trials due to non-safety related technical issues. Pre-meeting package for intercenter teleconference with CDER and CDRH representatives to discuss the device related aspects of the NDA. Verification of receipt of three copies of pre-meeting package; would like additional five copies. Response to request for five additional copies of SNO69 (dated May 11, 1998).
22-Apr-1998 29-Apr-1998 11-May-1998 15-May-1998 15-May-1998 22-May-1998	content/format of the NDA has been granted. Response to FDA fax regarding statistical comments on protocol C-94-057 and protocol FEN-USA-58. Notification of temporary suspension of clinical trials due to non-safety related technical issues. Pre-meeting package for intercenter teleconference with CDER and CDRH representatives to discuss the device related aspects of the NDA. Verification of receipt of three copies of pre-meeting package; would like additional five copies. Response to request for five additional copies of SN069 (dated May 11, 1998). Annual report covering the period of February 27, 1997 to February 26, 1998.
22-Apr-1998 29-Apr-1998 11-May-1998 15-May-1998	content/format of the NDA has been granted. Response to FDA fax regarding statistical comments on protocol C-94-057 and protocol FEN-USA-58. Notification of temporary suspension of clinical trials due to non-safety related technical issues. Pre-meeting package for intercenter teleconference with CDER and CDRH representatives to discuss the device related aspects of the NDA. Verification of receipt of three copies of pre-meeting package; would like additional five copies. Response to request for five additional copies of SN069 (dated May 11, 1998). Annual report covering the period of February 27, 1997 to February 26, 1998. Final Clinical Development Plan.
22-Apr-1998 29-Apr-1998 11-May-1998 15-May-1998 15-May-1998 22-May-1998	content/format of the NDA has been granted. Response to FDA fax regarding statistical comments on protocol C-94-057 and protocol FEN-USA-58. Notification of temporary suspension of clinical trials due to non-safety related technical issues. Pre-meeting package for intercenter teleconference with CDER and CDRH representatives to discuss the device related aspects of the NDA. Verification of receipt of three copies of pre-meeting package; would like additional five copies. Response to request for five additional copies of SN069 (dated May 11, 1998). Annual report covering the period of February 27, 1997 to February 26, 1998. Final Clinical Development Plan. Follow-up to query regarding the status of CDER/CDRH meeting
22-Apr-1998 29-Apr-1998 11-May-1998 15-May-1998 15-May-1998 22-May-1998 12-Jun-1998	content/format of the NDA has been granted. Response to FDA fax regarding statistical comments on protocol C-94-057 and protocol FEN-USA-58. Notification of temporary suspension of clinical trials due to non-safety related technical issues. Pre-meeting package for intercenter teleconference with CDER and CDRH representatives to discuss the device related aspects of the NDA. Verification of receipt of three copies of pre-meeting package; would like additional five copies. Response to request for five additional copies of SN069 (dated May 11, 1998). Annual report covering the period of February 27, 1997 to February 26, 1998. Final Clinical Development Plan.

Date of	
Contact	Summary of Contact
COILCACE	List of Janssen and ALZA attendees at the CDER/CDRH
10-Aug-1998	teleconference on August 6, 1998.
10 1149 1330	Sponsor's minutes from intercenter teleconference with
	CDER/CDRH representatives to discuss device related aspects
28-Aug-1998	of NDA.
	Meeting request to discuss previous agreements and
	communications with the Division in relation to the Clinical
28-Aug-1998	Development Plan.
	Protocol amendment - change in protocol and new investigator
23-Sep-1998	for the Phase I protocol C-94-060.
	Response to request for FDA meeting to confirm alignment of
	previous clinical agreements (request dated August 28, 1998,
6-Oct-1998	SN073).
	Regarding pending request to confirm previous clinical
26-Oct-1998	agreements still hold.
	Inquiry about availability of minutes from August 6, 1998
	CDER/CDRH/Janssen/ALZA meeting to discuss device
29-Oct-1998	documentation for NDA.
	Questions for Division for requested E-TRANS meeting per
4000	submission dated August 28, 1998 (SN073) and phone contact
10-Nov-1998	of October 22, 1998.
16-Nov-1998	Protocol amendment - new investigator for C-94-060.
	Background package for pending meeting request to discuss
	agreements/communications related to clinical development
18-Nov-1998	plan.
	Informing FDA that ALZA unable to attend clinical meeting in
1-Dec-1998	December per IND SN076.
	Voicemail requesting written withdrawal of meeting request
	since ALZA unable to attend a clinical meeting early
1-Dec-1998	December.
	Follow-up on October 19, 1998 fax regarding availability of
	minutes from August 6, 1998 intercenter meeting with
4-Dec-1998	CDER/CDRH/Janssen/ ALZA.
	Pre-meeting request to discuss previous agreements and communication with Division regarding clinical development
18-Dec-1998	plan.
10-060-1998	Verification that FDA received meeting request dated
	December 18, 1998 (SN077) for February meeting. Request for
11-Jan-1999	15 desk copies of background package.
	Follow-up to phone conversation tentatively setting meeting
	date of February 18, 1999 to discuss clinical development
18-Jan-1999	program.
	Acknowledgment of receipt of submission dated December 18,
	1998 (SN077). Confirmation of requested clinical development
13-Jan-1999	FDA meeting scheduled for February 18, 1999.
20-Jan-1999	Information amendment - CMC update.
	Follow-up of inquiries dated October 29, 1998 and December
	4, 1998 regarding availability of minutes from August 6,
20-Jan-1999_	1998 intercenter CDER/CDRH/Janssen/ALZA meeting.
	Desk copies of background package for pending meeting
	request to discuss previous agreements/communications with
20-Jan-1999	Division (SN076 and SN077).

Date of Contact	Summary of Contact
8-Feb-1999	Protocol amendment - new protocol C-94-067.
9-Feb-1999	Proposed agenda for February 18, 1999 meeting.
	Minutes from February 18, 1999 FDA meeting to discuss
18-Feb-1999	clinical program.
	Follow-up on fax dated October 29, 1998 and phone call dated
	December 4, 1998 regarding official FDA minutes from August
22-Feb-1999	6, 1998 CDER/CDRH/Janssen/ALZA meeting.
23-Feb-1999	Six additional copies of SN079 dated February 8, 1999.
	Request to meet with CDER and Division of Biopharmaceutics
26-Feb-1999	to discuss proposed finished product specifications.
	Information amendment - Pharmacology/Toxicology - final
10-Mar-1999	report for TR-97-1561-011 and TR-98-1561-031.
11-Mar-1999	Response to the meeting request filed on February 26, 1999 (SN080).
	FDA minutes from February 18, 1999 clinical development
17-Mar-1999	meeting.
	Background package for meeting with CDER and the Division of
29-Mar-1999	Biopharmaceutics on April 28, 1999.
	Requested changes to minutes for February 18, 1999 meeting;
19-Apr-1999	request for teleconference to discuss changes.
19-Apr-1999	Desk copies of SN083.
	1999 IND annual report covering the period of February 27,
23-Apr-1999	1998 to February 26, 1999.
-	Response to request to have team of FDA chemists come to
29-Apr-1999	ALZA to see the PSAL and SFTA.
_	Minutes from FDA meeting on April 28, 1999, including copies
	of summary overheads presented and agreed with FDA at close
30-Apr-1999	of meeting.
	Acknowledgment of receipt of April 19, 1999 correspondence
	(SNO83) requesting meeting to discuss response to Agency's
30-Apr-1999	February 18, 1999 meeting minutes.
	Inquiry regarding status of two outstanding items, one for
7-May-1999	JAN-1 and one for CPC-8.
10-May-1999	Recommendations regarding SN081 dated March 10, 1999.
27-May-1999	FDA minutes from April 28, 1999 JAN-1 Biopharm meeting.
	Inquiry regarding status of FDA minutes from August 1998
	teleconference (ref. SN072 dated August 28, 1998), FDA reply
	of CPC-8 response letter of March 5, 1998 (ref. SN019), FDA
	minutes from April 28, 1999 Biopharm meeting (ref. SN085
1-Jun-1999	dated April 30, 1999).
	FDA request for disk containing data related to April 28,
8-Jun-1999	1999 Biopharm meeting.
	Response to May 10, 1999 correspondence regarding inclusion
20 7 1000	of two nonclinical topical safety studies in the
30-Jun-1999	Investigator's Brochure.
	Response to FDA minutes from April 28, 1999 meeting with
	Division; submission of requested information related to
1 0 1000	claimed IVIVC; sponsor's request for written response to
1-Sep-1999	questions 4&5 from April 28, 1999 meeting.
22-Nov-1999	Confirmation that key agreements reached in prior CMC FDA

Date of	
Contact	Summary of Contact
	discussions still valid; sponsor minutes from September 25,
	1997 teleconference to discuss registration batch
	manufacturing and stability plans; request FDA minutes from
	August 6, 1998 teleconference
	Acknowledgment of receipt of CMC agreements submission sent
8-Dec-1999	November 22, 1999 (SN088).
	Voicemail in follow-up to message indicating FDA receipt of
21-Jan-2000	SNO88 (dated November 22, 1999)
	Request for FDA review of proposed change to the clinical
	development plan agreed upon at the ALZA/Janssen/FDA meeting
24-Feb-2000	on February 18, 1999.
9-Mar-2000	New supervisory Project Manager for Division
J Hai 2000	Record of phone communication with Supervisory Project
	Manager in relation to proposed change to clinical
14-Mar-2000	development program.
14-Mar-2000	Information amendment - pharmacology/toxicology final
	reports for nonclinical studies TR-99-1561-056 and TR-99-
27-Jun-2000	1562-057.
27-3411-2000	IND annual report covering the period of February 27, 1999
6-Jul-2000	to February 26, 2000.
0-041-2000	Request for preliminary review of new pharmacokinetic
18-Jul-2000	protocol C-2000-026.
18-341-2000	<u></u>
24 7 2000	Request for FDA response to proposal to submit clinical
24-Aug-2000	study synopsis for six terminated studies.
5-Sep-2000	Protocol amendment - new protocol C-94-067.
	Protocol amendment - new protocols, new investigators;
	information amendment - clinical, chemistry and microbiology
	for studies C-2000-005, C-2000-006, C-2000-007, C-2000-008,
11-Sep-2000	and C-2000-009.
	Response to ALZA request regarding status of FDA review of
	proposal to submit study synopsis for six safety and
19-Sep-2000	efficacy studies stopped in early 1998.
	Confirmation of FDA acceptance of proposal to submit
	clinical study synopses for six E-TRANS studies in the NDA
2-Oct-2000	(SN094 dated August 24, 2000).
	Protocol amendment - change in protocol C-94-067-04; new
5-Oct-2000	protocol C-2000-006-01; new investigator.
11 0-1 2000	
11-Oct-2000	Protocol amendment - new investigators.
12 0-+ 2000	Contacts between ALZA and FDA regarding status of C-94-067,
13-Oct-2000	ruled okay to proceed.
	Pharmacokinetic reviewer recommended to add 90 minute sample
00 0-1 0000	post start of first dose in C-2000-026 (SN093 dated July 18,
20-Oct-2000	2000).
20 0=+ 2000	Protocol amendment - change in protocol; information
30-Oct-2000	amendment - clinical for studies C-2000-026 and C-94-067.
	Table outlining the old Phase III study numbers, versus the
6 17 0000	new ALZA study numbers for the five Phase III protocols
6-Nov-2000	submitted to FDA on September 11, 2000 (SN096).
0000	Protocol amendment - new investigators for C-2000-005; C-
9-Nov-2000	2000-006; C-2000-007; and C-2000-009.
21-Nov-2000	Pre-NDA meeting request.
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Date of Contact	Summary of Contact
Contact	Summary of Contact
22-Nov-2000	Request for a January pre-NDA meeting.
	Request for a preliminary review of a new safety and
29-Nov-2000	clinical utility protocol (C-2000-030).
6-Dec-2000	Protocol amendment - new investigators for C-2000-005; C-2000-007; and C-2000-009.
7-Dec-2000	Response to meeting request dated November 21, 2000.
13-Dec-2000	Desk copy of previous SN103 (dated November 29, 2000).
15-Dec-2000	Background package for pre-NDA meeting.
15-Dec-2000	desk copies of background package for scheduled pre-NDA meeting and disk of cover letter and questions for FDA (SN105).
21-Dec-2000	Notification of termination of investigator participation in Protocol C-2000-007.
	Two comments from Medical Review Officer in relation to
22-Dec-2000	protocol C-2000-005.
5-Jan-2001	Protocol amendment - new investigators and revised 1572s for C-2000-005, C-2000-007, and C-2000-009.
	Response to Medical Review Officer comments on pediatric
5-Jan-2001	safety and efficacy protocol C-2000-005.
10-Jan-2001	Asking when and what serial number of the IND studies C-97-001; C-93-023; and C-94-067 were submitted.
11-Jan-2001	Protocol amendment - change in protocol C-2000-008.
12-Jan-2001	Informing that Anesthetics Division and CDRH reviewer had their internal prep. meeting on January 11, 2001 for our January 18, 2001 pre-NDA meeting and had questions.
16-Jan-2001	Response to FDA request for information - clinical.
20-Jan-2001	Response to request for pharmacokinetic simulations.
20 0411 2001	Informing that FDA has completed clinical review of SN098
1-Feb-2001	(dated October 5, 2000) and have comments.
	Informing that FDA has completed review of SN103 (dated
6-Feb-2001	November 29, 2000) and have comments/recommendations.
8-Feb-2001	Protocol amendment - new investigators for C-2000-007 and C-2000-009.
	Request for teleconference to discuss FDA's February 12, 2001 fax; request for clarification on intent of FDA's February 1, 2001 letter.
12-Feb-2001	:Follow-up to FDA's call on February 6
22-Feb-2001	Background for planned February 23, 2001 FDA teleconference to discuss February 1, 2001 FDA fax.
	Notification of termination of investigator participation in
22-Feb-2001	Protocol C-2000-009.
23-Feb-2001	ALZA minutes from February 23, 2001 FDA teleconference and IND table of clinical studies.
	Table of completed Phase 2 and Phase 3 studies, annotated to
	include reference to IND location of relevant safety data
23-Feb-2001	and final study reports for the short-stay surgery studies.
	Response to information requested at the February 23, 2001
26-Feb-2001	teleconference between ALZA and FDA.

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Date of	Summary of Contact
Contact	Response to messages from ALZA dated March 2, 2001 and March 5, 2001 regarding timing of receipt of FDA's minutes from January 18, 2001 pre-NDA meeting, and FDA input on the Home
5-Mar-2001	Safety protocol. Acknowledgment of receipt of call regarding IND SN113 (dated February 22, 2001). ALZA phone response to acknowledge
5-Mar-2001	voicemail. Interim safety data from ongoing Phase 3 clinical study of
7-Mar-2001	E-TRANS (fentanyl) in short-stay surgical patients. Request for information who is reviewing the Phase 2 data
7-Mar-2001	(C-96-020 and C-95-019) in the context of proposed protocol C-2000-030.
9-Mar-2001	Protocol amendment - new investigators for studies C-2000- 005 and C-2000-007. Requested clinical information related to complete Phase 2
12-Mar-2001	study C-96-020.
12-Mar-2001	Information related to Protocol C-2000-009 requested by FDA Request for information regarding notification of termination of Dr. Cork from participation in study C-2000-
12-Mar-2001	009 (IND SN113).
20-Mar-2001	FDA minutes of January 18, 2001 pre-NDA meeting. IND annual report covering the period of February 27, 2000
23-Mar-2001	to February 5, 2001.
30-Mar-2001	Information related to protocol C-2000-007
6-Apr-2001	Protocol amendment and information amendment for study C-2000-026.
16-Apr-2001	Type A meeting request. Acknowledgment of receipt of Type A meeting request (SN122 dated April 16, 2001) and indicating May 4, 2001 at 1pm only
18-Apr-2001	possible meeting date/time. Request for available monitoring reports for two terminated
19-Apr-2001	sites
25-Apr-2001	Contact regarding the April 16, 2001 Type A meeting request. Final question #1 and white paper related to previously
25-Apr-2001	submitted Type A meeting request.
2-May-2001	Protocol amendment - new investigators for study C-2000-005. Division's response to expanded Home Safety study questions
7-May-2001	(SN123 dated April 25, 2001). (Proposed agenda for May 10, 2001 FDA teleconference to
10-May-2001	discuss Home Safety Study. Protocol amendment - New protocol, new investigators -
14-May-2001	pediatric PK protocol C-2001-006 (in perioperative setting). Notification of the June 6, 2001 Type C FDA meeting
15-May-2001	(original letter dated May 1, 2001). Briefing package submitted for June 6, 2001 Type C FDA
21-May-2001	meeting. Package included ALZA's response to FDA's minutes of the January 18, 2001 pre-NDA meeting.
30-May-2001	FDA minutes of May 10, 2001 teleconference regarding home safety study (original letter dated May 16, 2001).
4-Jun-2001	Response to voicemail requesting pharmacokinetic data.

Contact Response to request for information pertaining to two clinical investigators who participated in the clinical program. Copy of overhead discussed with FDA at the close of the Jun 6, 2001 meeting. 2-Jul-2001 FDA meeting. 3-Jul-2001 FDA meeting. 6-Jun-2001 FDA meting. 6-Jun-2001 FDA meting. 6-Jun-2001 FDA minutes from June 6, 2001 meeting between ALZA and FDA. 5-Jul-2001 FDA minutes of the June 6, 2001 AP-22 meeting. Request for FDA review of proposed common name for the E-TRANS (fentanyl) acute system. 10-Aug-2001 Request for FDA review of proposed common name for the E-TRANS (fentanyl) acute system. 7-Sep-2001 Request for FDA review of protocol C-2001-011. 30-Aug-2001 Protocol amendment - new investigators for study C-2000-005 FDA comments on protocol C-2001-011 (submitted in SN131 dated August 13, 2001). Protocol amendment - New protocol and new investigator for protocol C-2001-009; Information amendment - clinical. Clinical information amendment submitted reflecting changes to 1572's previously submitted in SN126 with the original protocol C-2001-006. Regarding items related to the planned e-NDA and plan to fa in a proposal to submit the risk management plan to the NDA at the 6-7 month review period following NDA submission (SN130), which requested FDA review of proposed generic (common name) for E-TRANS (fentanyl) product. Submission of protocol amendment - new investigator; information amendment - clinical for studies C-2000-007, C-10-Jan-2002 2000-008, and C-2001-011. Response to inquiry regarding status of August 10, 2001 IND submission (SN130), which requested FDA review of proposed generic (common name) for E-TRANS (fentanyl) product. Submission of protocol amendment - change in protocol C-2001-011. General correspondence related to IND SN130 - request for review of proposed generic (common name) for the E-TRANS (fentanyl) product. Submission of protocol amendment - change in protocol C-2001-012. Protocol amendment - new investigator and information amendment - cl	<u> </u>	
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8-Aug-2002 investigator. 22-Aug-2002 Protocol amendment - change in protocol.	14-May-2002	
22-Aug-2002 Protocol amendment - change in protocol.	0.7	=
	0-Aug-2002	investigator.
	22-Aug-2002	Protocol amendment - change in protocol.
9-Oct-2002 Informing about the Clinical Trials Data Bank.		
	9-0ct-2002	Informing about the Clinical Trials Data Bank.

Date of	
Contact	Summary of Contact
	Request for a trademark consultation on the proposed
28-Apr-2003	Tradename
9-May-2003	Contact pertaining to the Tradename submission
J Hay 2000	IND annual report covering the period of February 27, 2002
9-May-2003	to February 26, 2003.
	Regarding meeting with Toni Nearing (WDC Liaison), Mark
	Kramer, and Patricia Love at the office of Combination
30-Jul-2003	Products on July 30, 2003.
	Minutes from teleconference held with the Office of Information Management Staff regarding planned electronic
:	NDA for E-TRANS (fentanyl); outline of e-NDA planned for
11-Aug-2003	submission in late September 2003.
	Protocol amendment - new protocol CAPSS-319 and new
	investigator information; information amendment - CMC
17-Dec-2003	information for protocol CAPSS-319.
06 R-h 000	Protocol Amendment - change in protocol CAPSS-319;
26-Feb-2004	Information Amendment - Change in contract packager Protocol Amendment - New Protocol CAPSS-320 and New
	Investigator Information; Information Amendment - CMC
2-Mar-2004	Information for Protocol CAPSS-320
	Dust and amount shows in market CARCC 220
2-Apr-2004	Protocol amendment - change in protocol CAPSS-320.
14-Apr-2004	Protocol Amendment: New Investigators
10 7 0004	Annual Report covering the reporting period of February 27,
19-Apr-2004	2003 to February 26, 2004.
14-May-2004	Protocol Amendment: New Investigators
26-May-2004	Protocol Amendment: New Investigators.
14-Jun-2004	Protocol Amendment: New Investigators.
23-Jun-2004	Protocol Amendment: New Investigators.
14-Jul-2004	Protocol Amendment: New Investigators.
21-Jul-2004	Protocol Amendment: New Investigators.
13-Aug-2004	Protocol Amendment: New Investigators.
18-Aug-2004	Protocol Amendment: New Investigators.
14-Sep-2004	Protocol Amendment: New Investigators.
15-Sep-2004	Protocol Amendment: New Investigators.
13-Oct-2004	Protocol Amendment: New Investigators.
13-Oct-2004	Protocol Amendment: New Investigators.
3-Nov-2004	Protocol Amendment: Change in Protocols CAPSS-319 and CAPSS-320
	Formal submission containing new investigator documentation
	for study CAPSS-319. This submission was assembled and sent
12-Nov-2004	by our CRO Pharmanet on behalf of ALZA.
	Formal submission containing new investigator documentation for study CAPSS-320. This submission was assembled and sent
12-Nov-2004	by our CRO Pharmanet on behalf of ALZA.
	Protocol Amendment: New Investigators.
16-Dec-2004	<u> </u>

Date of	
Contact	Summary of Contact
16-Dec-2004	Protocol Amendment: New Investigators.
7-Jan-2005	Protocol Amendment: Change in Protocol CAPSS-320.
14-Jan-2005	Protocol Amendment: New Investigators.
11-Feb-2005	Protocol Amendment: New Investigators.
14-Feb-2005	Protocol Amendment: Change in Protocol CAPSS-320.
11-Mar-2005	Protocol Amendment: New Investigators
31-Mar-2005	Annual Report for reporting period 2/27/2004 to 2/26/2005
15-Apr-2005	Protocol Amendment: New Investigators
2-Jun-2005	Protocol Amendment: New Protocol and New Investigator Info; Information Amendment
29-Jun-2005	Protocol Amendment: Change in Protocol C-2004-016 and New Investigator Information
25-Jul-2005	Protocol Amendment: New Investigators
25-Jul-2005	Amendments 3 & 4 of Protocol C-2004-016 (Serial No. 187) was submitted to the Agency on July 25, 2005.
2-Dec-2005	Protocol Amendment: New Investigators
17-Jan-2006	Protocol amendment - new protocol C-2005-028 and new investigator information; information amendment - CMC information for protocol C-2005-028 and supportive nonclinical data.
1/-Jan-2006	HOHCITHICAL GACA.
27-Jan-2006	Protocol amendment
24-Apr-2006	IND Annual Report 2/27/2005 to 2/26/2006

NDA Activities

Date	Summary of Contact
27-Oct-2000	Confirmation of NDA number (21-338) and the User Fee ID Number (4054).
12-Mar-2002	Request for a User Fee Identification Number for NDA 21-338 per instructions on FDA Form 3397 (User Fee Cover Sheet).
01.11	Response to ALZA's inquiry about setting up encrypted e-mail between ALZA and the Division to facilitate
21-Mar-2002	communication during review of the eNDA. Information on setting up encrypted e-mail between ALZA and the Division of Anesthetics, Critical Care, and
11-Apr-2002	Addiction Drug Products in preparation for the AP-22 NDA.
31 - Jan-2003	Inquiry about when ALZA plans to submit the NDA.
13-Jun-2003	Return telephone call regarding Dr. McNeil, Medical Reviewer who's been reviewing C-2002-027 in the IND and wondered about the status of the study. Informing of ALZA's plans to submit the e-NDA late September or early October 2003.
	DLT test generated by ALZA to confirm logging in process for official tape/submission. Levin forwarded e-mail to Ken Edmunds, electronic submissions coordinator.
3-Ju1-2003	Teleconference request for July 14, 2003 or July 28, 2003.
9-Jul-2003	Call to inform Compton of Regulatory Operations staff's July 7, 2003 e-mail to Levin to discuss aspects of the planned eNDA in late September or early October 2003.
29-Jul-2003	Teleconference to discuss the AP-22 eNDA test DLT tape submitted to CDER.
28-Aug-2003	User Fee sent by FedEx to the Mellon Client Service Center in Pittsburgh, PA.
	FDA advised to submit the DLT NDA tape, along with the originals of signed administrative documents in archival NDA jackets, and provided address for mailing of archival
23-Sep-2003	and desk copies. Submission of original new drug application (NDA) in
25-Sep-2003	electronic format. Regarding NDA filing date. 60 day filing date will be
10-0ct-2003	November 21, 2003. Looking to schedule filing meeting week of November 3, 2003.
	Acknowledgment of receipt of original NDA dated September 23, 2003. FDA internal filing meeting to take place week of November 23, 2003. Ten-month PDUFA review goal date is
15-Oct-2003	July 24, 2004. Request related to the AP-22 NDA case report forms for
6-Nov-2003	clinical studies C-94-057; C-94-058; and C-94-059. Response to November 6, 2003 phone request from MRO for
10-Nov-2003	case report form table of contents for clinical studies C-94-057; C-94-058; and C-94-059.
12-Nov-2003	Amendment - revised case report forms table of contents for clinical studies C-94-057, C-94-058, and C-94-059.

Date	Summary of Contact
Dace	Request from statistical reviewer for documentation
	related to data sets, and if possible, programs used to
13-Nov-2003	product efficacy results in reports and analysis.
13 100 2003	Amendment - CD-ROM containing combined sets of safety
	narratives organized by clinical/pharmacokinetics study
14-Nov-2003	and patient ID number.
14 110 2003	Asked if the request received via phone on November 13,
14-Nov-2003	2003 was a fileability issue request.
14 110 1 2003	Verification that the AP-22 NDA has been officially filed
25-Nov-2003	as of November 23, 2003.
23 1.01 2003	Amendment - CD-ROM containing documentation related to
	data sets and programs used to produce efficacy results in
3-Dec-2003	reports and analysis.
3 BCC 2003	NDA filing review completed (submissions dated September
	24, 2003; November 12, 2003; and November 14, 2003). No
	potential filing review issues noted to date. No major
	deficiencies noted thus far into the preliminary
5-Dec-2003	evaluation of the application.
	Regarding appropriate FDA contacts to discuss proposal for
17-Dec-2003	handling commercial product complaints.
	Request from a reviewing chemist to send placebo systems
2003-Dec-31 to	and any instructional materials needed to operate the
2004-Jan-02	system. ALZA response.
	Return call regarding possible meeting with FDA to discuss
	issues related to CDER/CDRH compliance grey zones eg
	complaints reporting, AE reporting, and jurisdictional
6-Jan-2004	issues for PAI.
	Message requesting information on the manufacturing flow
6-Jan-2004	and ALZA response.
20-Jan-2004	4-month cafety undate report
20-Jan-2004	4-month safety update report. Requested demonstrator systems of E-TRANS (fentanyl HCl)
21 700 2004	
21-Jan-2004	system.
23-Jan-2004	Question regarding ALZA's proposed risk management plan
23-Jan-2004	and ALZA response.
	Confirmation that AP-22 will not go to Advisory Committee, unless/until we decide to pursue an outpatient indication.
	Inquiry regarding when we will submit a more detailed risk
28-Jan-2004	management plan.
28-0a11-2004	Response to call from FDA dated January 28, 2004. FDA
	inquiry concerning timing of a more detailed risk
2-Feb-2004	management plan. being submitted to the NDA.
2 160 2004	Request for information regarding clinical section of the
12-Feb-2004	original NDA (dated September 23, 2003).
12 Teb 2004	Response to FDA's February 12, 2004 request for
12-Mar-2004	information
12 1101 2004	Request for FDA response regarding proposals for post-
	marketing safety reporting, product complaints
	investigation/reporting, and pre-approval inspectional
15-Mar-2004	jurisdiction.
10-Ma1-2004	Request for information regarding pharmacokinetic data
15-Mar-2004	sets for C-93-023-00 and C-2001-006-02
13-Mar-2004	
15-Mar-2004	Response to February 12, 2004 Information Request Letter.

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Date	Summary of Contact
21-Mar-2004	Response to FDA request for information
22-Mar-2004	Response to FDA's March 15, 2004 request for information.
	Request for review of proposed trade name IONSYS and
25-Mar-2004	generic product descriptor.
26-Mar-2004	Response to Project Manager's request for pharmacokinetic data (request dated March 15, 2004).
20 Mai 2004	Risk management proposal. Supercedes the risk management
2-Apr-2004	outline submitted with original eNDA.
8-Apr-2004	Requested Copy of Volume 1 of Electronic NDA 21-338
	Request for information regarding original NDA filing
	(dated September 23, 2003). CDRH has reviewed the device
12-Apr-2004	manufacturing section and requests additional information.
16-Apr-2004	Statistician would like to know if ALZA is planning to update the stability data and if so, when.
10 API 2004	NDA Amendment - Update of CMC Information; Revised
16-Apr-2004	Labeling.
	Copy of NDA Amendment dated April 16, 2004 was sent to FDA
16-Apr-2004	Field Office.
	Response to FDA question from statistician re: stability
19-Apr-2004	data.
19-Apr-2004	Confirmation of receipt of information re: stability data sent on April 16, 2004.
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22-Apr-2004	Copy of Clinical Information Request Letter.
22-Apr-2004 & 28-Apr-2004	Copy of CMC Information Request Letter.
30-Apr-2004	NDA Amendment - Response to Information Request Letter.
30-Apr-2004	Notification to District Office of NDA Amendment.
30 Hpz 2001	Response to Second Clinical Request for Information Letter
30-Apr-2004	for IONSYS
13-May-2004	Response to April 22, 2004 CMC Information Request Letter.
	Copy of cover letter for response to CMC Information
13-May-2004	Request letter sent to the Alameda District Office.
25-May-2004	Response to Request for Information.
nay 2001	Request for additional information following FDA monthly
	review meeting. FDA proposes a telecon for an information
25-May-2004	exchange (response to questions).
	Call to confirm June 15 telecon. Also, verification that
	the FDA wants both button pressing data and gel stability
27-May-2004	data from 9-mo.
	Confirmation of receipt of CMC/ device questions and notification that some of the items have been compiled and
	will be sent on 6/4. Also, notification that the
	technical group will be working to answer the remaining
3-Jun-2004	questions prior to the scheduled 6/15 telecon.
3-Jun-2004	Copy of CMC Information Request Letter.
	Requested Updated Stability Report; Response to Question
4-Jun-2004	Raised During Pre-approval Inspection.

Date	Summary of Contact
Date_	
	Follow-up on questions re: proposed tradename and post
	marketing complaints/ safety reporting. Also, follow-up
0.7	re: ALZA's request for clarification on question #7 of the
9-Jun-2004	most recent Information Request Letter.
	Request for FDA-input for the planned teleconference on
9-Jun-2004	June 15 following the internal FDA prep. Meeting.
11-Jun-2004	Response to Information Request Letter dated May 28, 2004.
11 0411 2001	Copies of cover letters for 6/4/04 Response to Request for
	9-month Corrective Action Lot Stability Update and
	Response to Question Raised during Pre-approval
	Inspection, and 6/11/04 Response to May 28, 2004
114 - Tun - 2004	_ =
14-Jun-2004	Information Request Letter.
14-Jun-2004	List of discussion topics for 6/15/04 teleconference.
	Copy of list of J&J attendees from the 6/15/04
16-Jun-2004	teleconference.
	Inquiry re: CMC Information Request Letter to be
21-Jun-2004	forthcoming from Agency
24-Jun-2004	Return call re: inquiry about the status of the CMC IRL.
29-Jun-2004 to	Response to points raised during teleconferences held on
30-Jun-2004	6/15/04 and 6/30/04.
00 0411 2001	Request for confirmation of receipt of response to points
	raised during 6/30 teleconference. Also, inquiry as to
	whether the primary review on the NDA are complete or if
1-Jul-2004	further questions might be forthcoming.
1-041-2004	Responses to Points Raised in the June 15, 2004 and June
1-Jul-2004	30, 2004 Teleconferences.
1 041 2001	Telecon to discuss reconciliation of the stability data
7-Jul-2004	and analysis from submissions dated 6/4/04 and 6/11/04.
7 041 2001	Request for clarification on a point re: adhesion
7-Jul-2004	discussed in the 7/7 teleconference.
7 041 2004	Copy of FDA minutes of the IONSYS CMC teleconference held
9-Jul-2004	on 7/7/04.
9 · 0 u I - 2 0 0 4	Teleconference and follow-up e-mail re: FDA question
0 7.1 2004	
9-Jul-2004	regarding the structure and chemical formula for R004380.
12 7-1 0004	Responses to Request for Methods Validation Document
13-Jul-2004	Package.
	Copies of cover letters for two electronic amendments for
	CMC submissions (18.1 and 19.1) forwarded to the FDA
16-Jul-2004	Alameda District Office.
	Response to CMC Memo dated 7/9/04; Formal Submission of
	Previously Communicated Nonclinical Response; Formal
17-Jul-2004	Submission of Revised Physician Insert.
,	Copy of NDA action letter (approvable) and CDRH Discipline
23-Jul-2004	Review letter received via e-mail.
	Hardcopy of NDA action letter (approvable) received from
	the Agency via regular mail. The CDRH Discipline Review
23-Jul-2004	letter was NOT received via hardcopy, but by e-mail only.
	Copy of NDA action letter (approvable) and CDRH Discipline
23-Jul-2004	Review letter received from the Agency via e-mail.
23-Jul-2004	Hardcopy of CDRH Discipline Review letter.
25-041-2004	maracopy or comm pracrieting mediem refret.

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Data	Summary of Contract
Date	Summary of Contact
28-Jul-2004	Request for clarification from the Agency regarding
thru 29-Jul-	expectations on response to the CDRH Discipline Review letter and the approvable letter. Also requested a
2004	teleconference with Dr. Rappaport.
2004	Returned call re: request for a telecon w/Dr. Rappaport.
	Jani stated that the response to the CDRH letter should be
	part of the complete response, but that issues raised in
29-Jul-2004	the CDRH letter were not approvability issues.
29-041-2004	Initial response to action letter as well as a request for
	a face-to-face meeting to clarify some of the points in
30-Jul-2004	the letter.
30-041-2004	Receipt of Action Letter and Intent to Amend the NDA with
30-Jul-2004	a Complete Response.
30-041 2004	Call from Compton to offer the date of September 10 for
	the AP-22 meeting to clarify issues in the NDA action
6-Aug-2004	letter.
O Aug 2004	Confirmation of Type A meeting with the Agency on
11-Aug-2004	September 10, 2004.
11 Aug 2004	Official letter from FDA granting Type A meeting to
11-Aug-2004	discuss approvable letter and CDRH letter.
11 Mag 2004	Copy of letter containing the Agency's comments on the
18-Aug-2004	proposed Risk Management Plan.
10 Aug 2004	Sponsor's questions for September 10, 2004 meeting w/FDA
24-Aug-2004	to discuss the approvable letter for NDA 21-338.
24 Mag 2004	Copy of slides presented by the FDA at the September 10
10-Sep-2004	Type A Post-Action Meeting.
10 BCP 2004	Questions re: when we can expect to receive the minutes of
	the 9/10 meeting, as well as inquiry as to whether it
21-Sep-2004	would be useful to send a video on how E-TRANS works.
31 30p 2001	Call re: desire to get an early review of the draft CDRH
27-Sep-2004	response.
30-Sep-2004	Questions re NDA 21-338.
7-Oct-2004	FDA review of IONSYS name.
8-Oct-2004	FDA minutes of the 9/10/04 IONSYS meeting.
	Type A Meeting Request: Clinical; Comments/Request to
	Correct Items in FDA's Minutes of September 10, 2004
29-Oct-2004	Meeting.
	Type A meeting request: Clinical; Comments/Request to
29-Oct-2004	correct items in FDA minutes of Sept 10, 2004 meeting
	Briefing Package for AP-22 Type A FDA Meeting (clinical)
15-Nov-2004	scheduled for December 2, 2004
	Notification briefing package for Dec 2 Type A meeting was
15-Nov-2004	Fed Ex'd
	Confirmation letter for Type A meeting request: clinical
17-Nov-2004	for IONSYS on December 2, 2004
	Cover letter and briefing package/question for FDA for Dec
17-Nov-2004	2, 2004 meeting
	Two Videos: Clinical Companion Video: Information for
	Healthcare Professionals and Clinical Companion:
17-Nov-2004	Information for Patients.
	Outcome of FDA's pre-meeting planned for December 2, 2004
23-Nov-2004	Type A meeting.

Date	Summary of Contact
Date	Agency Responses (Final) to sponsor's questions in meeting
9-Dec-2004	package for the IONSYS (Fentanyl HCl) product.
20-Dec-2004	Type B Meeting Request: CMC/CDRH Issues
4 to 5 Jan 2005	E-mail regarding potential CMC/CDRH meeting date
	Confirming Feb.10, 2005 CMC/CDRH meeting with the agency
6 to 7 Jan 2005	for NDA 21-338 Confirming of acceptance of Feb.10, 2005 CMC/CDRH meeting
6 to 7 Jan 2005	with the agency for NDA 21-338
13-Jan-2005	briefing package/questions for 2/10/05 FDA meeting
	FDA responses to questions posed for 2/10/05 CMC/CDRH
8-Feb-2005	meeting. FDA will pass on ALZA's clinical questions re:
9-Feb-2005	resubmission to medical officer
	ALZA's summary minutes of the Feb 10, 205 CMC/CDRH FDA
10-Feb-2005	teleconference.
25-Feb-2005	Clinical questions for the medical reviewer team leader at the FDA
	Formal submission of an e-mail sent to Kim Compton by ALZA
	on 2/25/05 containing clinical questions for the medical
1-Mar-2005	reviewer and/or medical team leader at the FDA FDA's minutes of the Feb. 10 teleconference IONSYS
	CDER/CDRH (CMC issues and draft response to CDRH
7-Mar-2005	Discipline review letter)
7-Mar-2005	E-TRANS Clinical questions
7 Hai 2005	sent response to FDA for E-TRANS fentanyl re: FDA
11-Mar-2005	questions on EU Trial Report FEN-PPA-401
23-Mar-2005	Follow up to Question 1 (adequacy of CDRH response)
31-Mar-2005	NDA 21-338 E-trans Fentanyl System - Device Issues
	Copy of planned TOC for the NDA resubmission was provided
8-Apr-2005	to the Agency for review.
	Agency confirmed its agreement on the Company's proposal to submit an abbreviated ICH study report for the EU trial
8-Apr-2005	FEN-PPA-401 in the NDA submission
	Type C Face-to-Face Meeting Request and Briefing Package:
	response to Office of Drug Safety Comments on the Original
22-Apr-2005	RiskMAP and revised draft risk minimization action plan.
	Informing that the Division will issue a letter to deny
	April 22, 2005 request for meeting to discuss the RiskMAP. The division felt the questions posed in the meeting
6-May-2005	request package could be addressed in written form.
2 130 3 2 0 0 0	Regarding the revised RiskMAP and ODS response that was
	submitted as pert of the meeting request (dated April 22,
9-May-2005	2005).
	Request for additional copies of the RiskMAP meeting
10-Marra 2005	package. Also included are responses to Kim's inquiry regarding the status of two items.
10-May-2005	Copy of ALZA's minutes of the April 1, 2005 teleconference
12-May-2005	submitted to the Agency.
	Sponsor's minutes of the April 1, 2005 teleconference with
12-May-2005	the Agency.

Date	Summary of Contact
	Letter advising that the Agency will provide written
1.6 14 0005	comments to questions in the proposed meeting request in
16-May-2005	lieu of a meeting. FDA's minutes of the April 1, 2005 Division/CDRH/OC
	teleconference. Purpose of the meeting was to discuss
27-May-2005	subject pertaining to the IONSYS NDA resubmission.
23-Jun-2005	Discussion of PK, AEs, and risk management plan
	Continuation of discussion regarding clinical questions for the medical reviewer [cf. RACRs dated 25-Feb-2005 and
23-Jun-2005	08-Apr-2005]
25 0uii 2005	voicemail with a question about where to find a referenced
20-Jul-2005	risk management analysis (D220005).
29-07-2005	RMP letter response to meeting request
29-07-2003	Call to thank the FDA for the esponses to our questions on
1-Aug-2005	the revised RiskMap.
	Call to discuss SPL and whether this is a requirement that
	would affect the IONSYS resubmission or is applicable only
8-Sep-2005	to new registration applications made after Oct. 31, 2005.
21-Sep-2005	Follow-up to the 9/8/05 conversation re: SPL requirement
	Courtesy message informing Kim Compton of ALZA's plan for
	submitting the resubmission/complete response on 11/21/05
	and the content and format in which the submission will be
16-Nov-2005	sent to the July 2004 approvable letter.
18-Nov-2005	confirmed FDA mailing address & Agency has 14 days to determine if response is complete.
	determine in response is complete.
21-Nov-2005	Submission of Complete Response
	Request by the Agency to submit requested information on
22-Nov-2005	the specification no later than March 15, 2006 to facilitate the review process.
22-NOV-2003	Confirmation that the Agency received ALZA's response to
	the July 23, 2004 Action Letter. The Agency will decide
	by December 6, 2005 whether it is a complete response and
29-Nov-2005	thus restarts the clock.
	Follow-up to FDA's inquiry about submission of information
1-Dec-2005	related to specification.
6-Dec-2005 to	Notification the NDA resubmission submitted on 11/11/05 is
7-Dec-2005	a complete response to FDA's 7/23/05 approvable letter. Official letter from the FDA re: the IONSYS NDA
	resubmission. The Agency considers the resubmission as a
	complete, Class 2 response to the July 23, 2004 action
9-Dec-2005	letter and the PDUFA user fee goal date is May 22, 2006.
13-Dec-2005	Voicemail regarding analytical lab
13-Dec-2005	Voicemail regarding one of J&J sites
6-Jan-2006	Withdrawal of Analytical Testing Laboratory.
	E-mail to inform FDA project manager of the IONSYS EMEA
	communication regarding the delay in EU launch due to a
	recently identified issue with the commercial
17-Jan-2006	manufacturing process for IONSYS.

Date	Summary of Contact
	Request from the Agency for samples (placebo) of the
2-Feb-2006	IONSYS system.
	Samples of IONSYS System (demo units without gels) sent to
10-Feb-2006	Agency for review.
	FDA rec'd the IONSYS systems (20 sample units) ALZA
14-Feb-2006	submitted 2/10/06
	Request from the Agency for a brief teleconference for
23-Feb-2006 to	February 27, 2006 to clarify a couple of items on the
24-Feb-2006	proposed RMP for IONSYS.
	Telecon at FDA's request to discuss some items related to
	revised Risk Map (submitted April 2005) and NDA
07 7 1 0006	resubmission response to FDA's July 29, 2005 letter with
27-Feb-2006	comments on the revised Risk Map.
	CMC question for the Agency regarding the difference in
10 Man 2006	POC and CAL lots. Heads up that comments/requests from
10-Mar-2006	CDRH will be coming soon. NDA amendment containing revised specs for Impurity A, B
14-Mar-2006	and FC1003
14-Ma1-2000	Field copy of NDA amendment containing revise specs for
15-Mar-2006	Impurity A, B and FC1003 sent to District Office
21-Mar-2006	Submitted the revised IONSYS RiskMAP (March 2006 edition)
	Questions from the Center of Devices and Radiological
22-Mar-2006	Health (CDRH)
24-Mar-2006	FDA teleconference scheduled for March 30, 2006
	List of FDA invited attendees for the 3/30/06
	teleconference to discuss items outlined in the 3/22/06
27-Mar-2006 to	CDRH letter. Also, FDA comments on the proposed patients
28-Mar-2006	instructions for use for IONSYS.
	List of ALZA/J&J attendees and questions for the Agency
	for the March 30, 2006 teleconference to discuss comments
29-Mar-2006	from the March 22, 2006 CDRH letter.
	Agency is not planning to issue minutes for the March 30,
	2006 teleconference. Company targeting to submit a bulk of
	the responses to the March 22, 2006 CDRH letter to the
30-Mar-2006	Agency by April 7th and the remaining responses by April 14th.
30-Ma1-2006	Follow-up action item from telecon w/ FDA re: educational
	materials and proposed revisions to FDA's version of PI
4-Apr-2006	labeling
4-Apr-2006	Telecon at FDA's request regarding Risk Map/review issues
5-Apr-2006_	Response to request for information
6-Apr-2006	Response to 3/22/06 information request letter
	Educational materials sent electronically as a follow-up
12-Apr-2006	to 4/11/06 email request
	Posponse to proposed educational material request
13-Apr-2006	Response to proposed educational material request Response to 3/22/06 information request letter (remaining
13-Apr-2006	questions)
13 Apr 2000	Follow-up to the April 3, 2006 telecon: email the EU PI
	and PPI and submit in writing where ALZA stands on the
19-Apr-2006	setting for IONSYS.
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Date	Summary of Contact
	ALZA's response to Ms. Compton's email 4/18/06 with a
19-Apr-2006	request to send European PI (SmPC) and PPI for IONSYS.
	Request from FDA (CMC team) to update the drug product
20 7 2006	specifications table to include the proposed test and
20-Apr-2006	acceptance criteria for "Dose Charge." ALZA's response to the issue at the April, 3 2006 FDA
	teleconference, regarding the appropriate setting for use
21-Apr-2006	of IONSYS.
22 142 200	Re: IONSYS and Duragesic, tcon and packaging
21-Apr-2006	consideration.
	Response to additional requests received via e-mail from
24-Apr-2006	FDA on April 18, 2006. (vn 38)
	List of participants in ALZA/FDA teleconference re: IONSYS
24-Apr-2006	NDA.
	Response to the April 20, 2006 request from CMC reviewer
0000	to add dose charge to the drug product specification
24-Apr-2006	(response to Question 4 in the July 9, 2004 amendment).
24-Apr-2006	Tcon to discuss disposal and packaging label issue
27-Apr-2006	Proposed revised label text re: red tab on IONSYS
	Clinical response in follow up to the April 24, 2006
30-Apr-2006	teleconference with FDA.
	Response to two information requests from the April 24,
	2006 telecon regarding: 1) pulling on the red tab of the
	IONSYS system and 2) clarification supporting nurses'
1-May-2006	understanding of the appropriate use of IONSYS.
1 1 1 2006	Acknowledgment of receipt of ALZA's clinical response, in
1-May-2006	follow up to the April 24, 2006 telecon with FDA. CMC request and to send validation pkg for SFTA test
2-May-2006	method
5-May-2006	Questions from FDA re: labeling items
10-May-2006	Response to FDA Request for SFTA Method Validation Packag
10-May-2006	Response to FDA 5/5/05 letter re: labeling items
	FDA acknowledgement re: Patient Bedside Sheet w/ no action
11-May-2006	needed from ALZA
11-May-2006	PT Bedside Sheet - one small change
11-May-2006	Telephone call pertaining to IONSYS labeling
11-May-2006	Response to May 5, 2006 FDA request
11-May-2006	Provided revised drug product specs to FDA
12-May-2006	FDA's comments on the IONSYS labeling (PI).
	Letter from the FDA containing Office of Drug
10 11 0005	Safety/Controlled Substances Staff (ODS/CSS) comments on
12-May-2006	the IONSYS RiskMAP.
15-May-2006	Comments from FDA requesting revisions to the carton and container labels.
15-May-2006	Physician's labeling and Word version of the patient
16-May-2006	labeling for IONSYS sent via e-mail to FDA.
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Date	Summary of Contact
Date	Response to May 12, 2006 and May 16, 2006 FDA request for
	IONSYS labeling (PI and patient bedside information
17-May-2006	sheet).
17-May-2006	
	FDA Project Manager confirmed receipt of the Physician's
17 1/ 0006	labeling and Word version of the patient labeling that was
17-May-2006	sent on a CD via FedEx to FDA.
	Modification to section of the PI, as discussed during the
18-May-2006	May 18, 2006 telecon w/FDA.
18-May-2006	IONSYS Physicial label (PI) from the FDA.
	FDA Project Manager confirmed receipt of the IONSYS May
18-May-2006	18, 2006 submission.
	Revised color mock up of the IONSYS system print.
	"Patient-activated" has been deleted and replaced with "40
18-May-2006	mcg/activation" per FDA request.
-	Response to FDA Discipline Review letter fromODS/CSS dated
18-May-2006	May 12, 2006 regarding the IONSYS RiskMAP.
	Response to FDA Discipline Review Letter dated May 12,
18-May-2006	2006 on the IONSYS RiskMAP.
	Submission related to the Proposed Draft Label (PI) for
	IONSYS. The cover letter is dated May 19, 2006 and the
	electronic version of this submission was sent via secure
19-May-2006	e-mail to FDA on May 19, 2006.
	Response to FDA inquiry re: exclusivity. In the cover
	letter of the original NDA, ALZA requested 3 years
	exclusivity due to the fact that we conducted significant
19-May-2006	clinical trials for IONSYS (per CFR 21 314.108).
	Submission of the subanalysis for the Nurse Ease of Care
	(EOC). Information was sent on May 18, 2006 via secure e-
19-May-2006	mail to FDA.
-	Confirmation from FDA regarding acknowledgement of minor
19-May-2006	typo/correction to p.18 of the PI.
	ALZA's final draft of the IONSYS label submitted to the
19-May-2006	Agency.
-	Followup on the subject of agreement with the Agency on
	the Risk Management Plan. Request to come to an agreement
	on a reasonable timeframe in which agreement on the Risk
19-May-2006	Management Plan would be obtained with the Agency.
	Dialogue between the Sponsor and the Agency regarding the
	timeframe related to agreement on the Risk Management Plan
19-May-2006	for IONSYS with the Agency.
21-May-2006	Discussion related to Risk Management Plan (RiskMAP).
22-May-2006	Approval letter for IONSYS
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